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Table 3. ADVANTAGE: Prior medication related to the cardiovascular system

	Rofecoxib 25 mg/d N= 2785		Naproxen 1000 mg/d N= 2772	
	n	%	n	%
Renin-angiotensin system	593	(21.3)	614	(22.2)
Antihypertensives	132	(4.7)	149	(5.4)
Beta blocking	390	(14.0)	418	(15.1)
Calcium channel blockers	408	(14.6)	440	(15.9)
Cardiac therapy	114	(4.1)	127	(4.6)
Diuretics	536	(19.2)	560	(20.0)
Peripheral vasodilators	13	(0.5)	16	(0.6)
Serum lipid reducing agents	564	(20.3)	523	(18.9)

(source Sponsor's Table 14, appendix 1.4)

Table 4. Prior use of antithrombotic agents

	ROFECOXIB 25 MG N= 2785		NAPROXEN 1000 MG N= 2772	
	n	%	n	%
Clopidogrel bisulfate	4	(0.1)	7	(0.3)
Dipyridamole	4	(0.1)	10	(0.4)
Ticlopidine	1	(0.1)	0	(0.0)
Warfarin sodium	6	(0.3)	1	(0.0)

(source Sponsor's Table 14, appendix 1.4)

Reviewer's comment: Similar percentage of patients had received cardiovascular medication and discontinued antithrombotic therapy within 30 days of entering the study. At the reviewer's request the sponsor provided information that none of these patients developed a serious cardiovascular thrombotic event during the trial.

Table 5. Advantage: Prior use of hormonal therapy

	Rofecoxib 25 mg/d N= 2785		Naproxen 1000 mg/d N= 2772	
	n	%	n	%
Endocrine therapy ¹	86	(3.1)	64	(2.3)
Sex hormones and modulators of the genital system ²	875	(31.4)	859	(31.0)

Source: Sponsor's Table 14, Appendix 4.1) ¹: Endocrine therapy includes mainly raloxifene and tamoxifen. ²: Sex hormones includes mainly estrogenic and progesterone-related hormones.

There were no significant differences in the number of patients who took hormone replacement therapy (HRT) prior to the trial. Approximately 31 % of patients took "hormones and modulators of the genital system" during the trial, including different estrogenic preparations with or without progesterone (n=741 and 720) progestins alone (n=145 and 151) or testosterone (n=12 and 10) in the rofecoxib and naproxen group respectively.

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ASA was a prior medication in 470 (16.9%) and 474 (17.1%) patients in the rofecoxib and naproxen groups, respectively. Aspirin is listed among the analgesic, regardless of the dose that was taken.

Reviewer's comment. The sponsor did not specify the dose of ASA taken prior to entry. Of note, 360 and 372 patients used concomitant ASA during the study, in the rofecoxib and naproxen group, respectively. Therefore, approximately 100 patients in each arm discontinued the use of ASA before entering the study. At the reviewer's request the sponsor provided information that only one of the patients who discontinued ASA prior to study entry developed a serious CV thrombotic event (AN 2401) who was taking ASA 1300 mg/day for the treatment of OA, not for cardioprotection. This patient (on rofecoxib 25 mg) developed superficial venous thrombophlebitis. The event was not confirmed as serious cardiovascular thrombotic by the adjudication committee.

1.4.5 Exposure

Although designed as a 3-month study, actual exposure was significantly shorter. Median exposure to both rofecoxib and naproxen was 84 days (mean was approximately 69 ± 30 days). Overall, most of the patients were compliant with dosing (88.8%). The percentage of patients with 80% compliance was similar in both treatment groups.

1.5 Safety Results

1.5.1 Overall safety

There was no overall advantage for rofecoxib 25 mg/d over naproxen 1000 mg/d. The numbers of patients with one or more adverse events (AE's), serious AE's, who died or discontinued due to an AE were similar in both treatment groups.

Table 6. NDA 21-042. ADVANTAGE study. Clinical Adverse Event Summary.

	Rofecoxib 25 mg/d (N= 2785)		Naproxen 500 mg b. (N= 2772)	
With one or more AEs	1814	(65.1)	1825	(65.8)
With serious AEs	68	(2.4)	72	(2.6)
Who died	5	(0.2)	4	(0.1)
Who discontinued due to an AE	374	(13.4)	386	(13.9)

Source sponsor's table 21.

1.5.1.1 Deaths

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There were 5 deaths in the rofecoxib group (0.2%) and 4 in the naproxen group (0.1%) (Table 7). Except for one 56-year-old patient who died of complications of gallbladder carcinoma, all patients were older than 70 years.

Four of the five deaths in the rofecoxib group were due to cardiovascular causes (three sudden death, one ruptured aortic aneurism). There were no cardiovascular deaths in the naproxen group. Of note, one patient with prior borderline renal function taking furosemide died of complications of acute renal failure in the naproxen group. None of the deaths were considered by the investigator to be drug-related. Narratives of deaths are presented in Appendix 1.

Table 7. Listing of Deaths in the ADVANTAGE study

AN/site	Age/ sex	Prior Medical History	Concomitant medications	Cause of death	Day #
Rofecoxib					
5005/065	73 F	HTN, lip, K	none	Sudden death	40
3700/200	74 M	DM, CAD, CABG, lip	ASA, ACE(-), statin, glybur	Sudden death	42
4856/210	71 F	-	none	Astrocytoma	120
4049/658	71 M	A fib, HTN, CAD,	digoxin, diltiazem	Sudden death	60
3423/679	75 F	Hematuria	none	Rupture aortic aneur	42
Naproxen					
1841/059	56 F	-	none	Gallbladder Ca.	120
7154/702	79 F	DM, HTN, anasarca	aleandronate, glypizide	Pancreatic Ca.	90
3105/777	74 F	HTN,CHF,depression, hyperuricemia, Cr. 1.5	verapamil, ACE(-), lasix, fluoxetine	Acute renal failure	40
7114/831	78 M	COPD, PVD, smoker	ASA	Lung Ca	60

Source: Advantage CSR.

Reviewer's comment:

Of note, the cause of death for patient # 065 5005 (on rofecoxib), had been listed by the investigator as hypertensive heart disease and not referred for adjudication as a potential cardiovascular thrombotic event to the cardiovascular adjudication committee. The patient called her son complaining of chest pain and by the time the son arrived she was dead. In the opinion of this medical reviewer, the cause of death for this patient was sudden death, which would in fact meet criteria for cardiovascular thrombotic event.

1.5.1.2 Serious AEs

A total of 140 patients - 68 (2.4%) in the rofecoxib group and 72 (2.6%) in the naproxen group - had at least one serious clinical AE during the study.

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The findings of this large but relatively short study (3 months) with the rofecoxib 25 mg dose are consistent with those in the longer term (9-month) VIGOR study at the 50 mg dose. Again there is a lower number of digestive system related AE's in the rofecoxib group, but there is no overall advantage of rofecoxib over naproxen.

Table 8. Serious AE's occurring in two or more patients in ADVANTAGE.

	Rofecoxib 25 mg	Naproxen 1000 mg
	N= 2785 n %	N= 2772 n %
Patients with at least one serious AE	68 (2.4)	72 (2.6)
Body as a whole	7 (0.3)	10 (0.4)
Cardiovascular system	23 (0.8)	17 (0.6)
Digestive system	7 (0.3)	21 (0.8)
Endocrine	1 (0.0)	1 (0.0)
Hemic and lymphatic	1 (<0.1)	2 (0.1)
Hepatobiliary system	3 (0.1)	1 (<0.1)
Musculoskeletal system	7 (0.3)	7 (0.3)
Nervous system	4 (0.1)	2 (0.1)
Psychiatric disorder	4 (0.1)	1 (0.0)
Respiratory system	6 (0.2)	5 (0.2)

(Source sponsor's table 24).

Reviewer's comment: Of note, the dose of rofecoxib used in this trial is half of the dose used in VIGOR but the dose of naproxen is the same (500 mg bid) in both trials. The incidence of serious adverse events in ADVANTAGE (2.4 and 2.6% in rofecoxib and naproxen respectively) is much lower than in VIGOR (9.3 and 7.8% in rofecoxib and naproxen respectively). This observation may be in part explained by the shorter duration of the study and the different population (OA in ADVANTAGE, RA in VIGOR).

A table of serious events that required hospitalizations is presented in Appendix 2. Again, there was no overall advantage of rofecoxib over naproxen. Twice the number of patients required hospitalization for GI related serious AE's in the naproxen group (5 and 12 for rofecoxib and naproxen respectively) and a numerically higher number of patients required hospitalization for CV related events in the rofecoxib group (18 and 13 on rofecoxib and naproxen respectively).

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1.5.1.3 Serious AE's by ASA use

Table 9. ADVANTAGE study. Concomitant ASA use during study by dose*

	ROFECOXIB 25 MG N= 2785		NAPROXEN 1000 MG N= 2772	
	n	%	n	%
Low dose (20-325 mg/day)	352	(12.7)	367	(13.3)
Non-ASA user (<20 mg/day)	2425	(87.0)	2400	(86.5)
Other ASA user (>325 mg/day)	8	(0.3)	5	(0.2)

* Categories defined by the sponsor for this study as follows: "Low dose": 20 to 325 mg/day; "Non user": Average of <20 mg/day or following a CV event; "Other": >325 mg/day or started therapy during study. Source: Table 10 Advantage CSR and response to request for information submitted by sponsor 6/29/01.

Distribution of ASA use in the ADVANTAGE study is presented in Table 8. Analysis of serious AE's by ASA use demonstrated a similar incidence of events in the low aspirin users and non-users, except for the cardiovascular and the digestive system.

Table 10. Serious AE's by ASA use (events with incidence 0.5%), as reported by investigators.

	Rofecoxib N= 2785		Naproxen N= 2772	
	Non ASA users	Low dose ASA	Non ASA users	Low dose ASA
	N= 2422	N= 352	N= 2398	N= 367
	n (%)	n (%)	n (%)	n (%)
Patients with at least one event in any body system	54 (2.2)	14 (4.0)	59 (2.5)	12 (3.3)
Cardiovascular	16 (0.7)	7 (2.0)	— (0.6)	2 (0.5)
MI	3	2	1	-
Sudden death	2	1	-	-
Digestive system	5 (0.2)	2 (0.6)	18 (0.8)	3 (0.8)

Source: Corrected Table 68 of Advantage CSR submitted 8/801).

Reviewer's comment:

In this short trial, the data suggest that co-use of low dose ASA increased the risk of serious GI events for rofecoxib (0.6%) but not for naproxen (0.8%, unchanged). The trial, however, was not designed to adequately assess the long-term effect of co-administration of ASA in the gastrointestinal system.

Most importantly, patients with known cardiovascular risk —as defined by patients using low dose prophylactic ASA - receiving rofecoxib had four fold

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more serious cardiovascular adverse events than those receiving naproxen (2.0% vs 0.5%).

In this three-month study with a dose of rofecoxib approved for chronic use in patients with OA, those on cardiovascular prophylaxis showed a trend towards more cardiovascular events (2.0 %) than those not on aspirin in the rofecoxib group (0.7%). Presumably, these patients are at higher cardiovascular risk than those not taking aspirin, therefore the finding is not unexpected. However, this was not the case in the naproxen treatment group (0.5 % and 0.6 %, for those who were and were not on ASA, respectively).

The sponsor has speculated that the excess risk of cardiovascular thrombotic events in the rofecoxib group as compared to naproxen in the VIGOR study may be due to the lack of anti-platelet effect of rofecoxib and has proposed that addition of low dose ASA in high risk patients may prevent the problem. The limited data on rofecoxib and ASA use from the ADVANTAGE study suggest that low dose ASA in patients with prior cardiovascular history, might not eliminate the excess risk of serious cardiovascular events of rofecoxib compared to naproxen.

1.5.1.3 Dropouts due to adverse events

Table 11. ADVANTAGE. Dropouts due to adverse events

	Rofecoxib N= 2785		Naproxen N= 2772	
	n	%	n	%
Number of AE dropouts	374	(13.4)	386	(13.9)
Body as a whole	87	(3.1)	102	(3.7)
Cardiovascular	40	(1.4)	21	(0.8)
Digestive system	113	(4.1)	142	(5.1)
Endocrine	1	(0.0)	0	(0.0)
Eyes, ears, nose and throat	7	(0.3)	8	(0.3)
Hemic and lymphatic	1	(0.0)	2	(0.1)
Hepatobiliar system	2	(0.1)	1	(0.0)
Musculoskeletal system	45	(1.6)	49	(1.8)
Nervous system	30	(1.1)	24	(0.9)
Psychiatric disorder	14	(0.5)	7	(0.3)
Respiratory system	8	(0.3)	10	(0.4)
Skin and skin appendages	21	(0.8)	23	(0.8)

Reviewer's comment:

Consistent with the VIGOR study, there was no overall advantage of rofecoxib 25 mg/day over naproxen 1000 mg/day, based on the number of dropouts due

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to AE's. The percentages of dropouts due to AE's in the ADVANTAGE study are similar to those in VIGOR (15.9 and 15.8% for rofecoxib and naproxen, respectively). Of note, the number of dropouts due to cardiovascular related events was almost twice in the rofecoxib arm (n=40, 1.4%) when compared to the naproxen arm (n=21, 0.8%).

1.5.1.4 Most common adverse events

The total number of adverse experiences was approximately 65% in each treatment group. Adverse experiences by body system were generally also similar between treatment groups, including the digestive system (24% and 26 % in the rofecoxib and naproxen arms, respectively). Again there was no overall advantage of rofecoxib 25 mg/day over naproxen 1000 mg /day.

1.5.1.5 Laboratory adverse events

Mean changes

Laboratory measurements were taken at baseline and at Week 12. Mean changes from baseline in each laboratory parameter were small and comparable between treatment groups, including hemoglobin, hematocrit, BUN, creatinine and liver function tests.

Serious Laboratory Adverse Experiences

Only one patient had a serious laboratory AE during the study. The event was moderately decreased hemoglobin. No action was taken with regard to study drug.

Discontinuations Due to Laboratory Adverse Experiences

Eleven patients (0.4%) in the rofecoxib group and 6 (0.2%) in the naproxen group, were withdrawn from the study due to a laboratory AE. None of them were serious.

1.5.1.6 Vital signs

Vital signs (diastolic and systolic blood pressure, heart rate, and respiration rate) were measured at each study visit. Mean changes and exceeding limits of change from baseline at week 6 and 12 were analyzed.

Maximum increases in systolic blood pressure at Week 12 were 94 mmHg and 60 mmHg for the rofecoxib and naproxen groups, respectively; **mean increases** were 1.04 mmHg and 0 mmHg in the rofecoxib and naproxen groups respectively. Maximum increases in diastolic blood pressure at Week 12 were 40 mmHg and 30 mmHg in the rofecoxib and naproxen treatment groups respectively; **mean increases** were 0.32 mmHg and -0.66 mmHg in the rofecoxib and naproxen groups, respectively.

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Defined limits of change for blood pressure were as follows: systolic blood pressure >140 mmHg with an increase from baseline >20 mmHg at either week 6 or 12; and diastolic blood pressure >90 mmHg with an increase from baseline >15 mmHg at either week 6 or 12.

Percentage of patients who exceeded the limit change for SBP at either week 6 or 12 were 10.7% and 9.1% for rofecoxib and naproxen, respectively. The percentage of patients who exceeded limits in SBP at both week 6 and 12 were 2.3% and 1.6% in the rofecoxib and naproxen groups, respectively. Few patients in either treatment group (0.5%) had a change in diastolic blood pressure exceeding the defined limit at both visits.

Reviewer's comment: Patients in the rofecoxib group tended to have a larger increase in blood pressure compared to the naproxen group, although, in this three-month study, the differences in blood pressure changes were small.

1.5.2 Analyses of Cardiovascular Safety

1.5.2.1 Cardiovascular thrombotic events. Serious cardiovascular (CV) AEs occurring in a patient while on study treatment or within 14 days of discontinuation of study treatment were reviewed by the sponsor for inclusion in the adjudication process.

Of the 23 and 17 investigator reported serious CV AEs in the rofecoxib and naproxen arm respectively, 14 and 13 were considered by the sponsor to be thrombotic-related (as per Merck's Vascular SAE Terms Eligible for Case Adjudication, Appendix 3) and referred for blinded adjudication to the Cardiovascular Adjudication Committee. Two additional cases obtained through the WAES (Worldwide Adverse Event System) database were also referred for adjudication by the sponsor, making a total of **14 and 15 cases referred** for adjudication from the rofecoxib and naproxen arms, respectively.

Reviewer's comment: The two cases referred for adjudication from the WAES database were not technically investigator reported events. It is unclear whether these cases represent unblinded data (both cases were on naproxen).

Investigator reported serious CV AEs are presented in the following table.

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Table 12. Investigator reported Serious AE's related to the CV system by ASA use

	Rofecoxib			Naproxen		
	All N= 2785	No ASA N= 2425	ASA N=352	All N= 2772	No ASA N= 2400	ASA N= 367
	n %	events	events	n %	events	events
Any CV related event	23 (0.8)	16 (0.7)	7 (2.0)	17 (0.6)	15 (0.6)	2 (0.5)
Acute Myocardial Infarction	1	-	1	-	-	-
Arterial Occlusion	1	1	-	-	-	-
Arterial rupture	1	1	-	-	-	-
Atherosclerosis	1	1	-	1	1	-
Atrial fibrillation	2	2	-	1	-	1
Cardiac arrest	-	-	-	1	1	-
Cardiovascular disorder	1	-	1	-	-	-
Carotid artery obstruction	1	-	1	-	-	-
Cerebellar hemorrhage	-	-	-	1	1	-
Cerebral aneurysm	1	1	-	-	-	-
Cerebral infarction	-	-	-	1	1	-
Cerebrovascular accident	-	-	-	3	3	0
Congestive heart failure	4	3	1	2	2	-
Coronary artery disease	2	1	1	2	1	1
Deep venous thrombosis	-	-	-	3	3	-
Hypertension	1	1	-	-	-	-
HTN heart disease	1	1	-	-	-	-
Myocardial infarction	3	2	1	1	1	-
MI- age indetermined	1	-	1	-	-	-
Non-Q wave MI	1	1	-	-	-	-
Pulmonary edema	1	1	-	-	-	-
Sick sinus syndrome	1	1	-	1	1	-
Subarachnoid hemorrhage	1	1	-	-	-	-
Supraventricular tachycardia	1	1	-	1	1	-
Third degree AV block	1	-	1	-	-	-
Thrombophlebitis	1	-	1	-	-	-
Transient ischemic attach	2	2	-	2	2	-
Unstable angina	1	1	-	-	-	-
Vasospasm	1	1	-	-	-	-
Ventricular fibrillation	-	-	-	1	1	-
Ventricular tachycardia	1	-	1	-	-	-

(Source: sponsor's tables 60, 68 & 69, corrected tables submitted 8/8/01). (N= patients randomized; n= patients with events). This list does not include two additional cases obtained though WAES.

Reviewer's comment: The incidence of investigator related serious CV AEs was higher among those patients taking concomitant aspirin in the rofecoxib group. The subgroup of patients taking aspirin presented four fold more serious CV adverse events in the rofecoxib group than the naproxen group.

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1.5.2.2 Adjudicated serious CV/thrombotic events.

Each clinical event referred for adjudication was reviewed blindly by three cardiologists (CV adjudication committee). The criteria for adjudication of cardiovascular serious thrombotic events were the same as the ones used in the VIGOR study.

Table 13. Criteria for Adjudication of Cardiovascular Serious Thrombotic events

a. Coronary – <i>Cardiology</i>
1. Acute MI (fatal or non-fatal)
a. Spontaneous
b. Secondary to an antecedent stressor (major surgery, GI bleed)
c. Complication of PTCA or coronary revascularization procedure
2. Unstable angina pectoris
3. Cardiac (atrial or ventricular thrombus)
4. Resuscitated cardiac arrest (without identified cause listed elsewhere)
5. Sudden or unexplained death
b. Peripheral (other than cardiac or cerebrovascular) vascular – <i>Peripheral Vascular</i>
1. Pulmonary embolism (fatal or non-fatal)
a. Spontaneous
b. Secondary to an antecedent stressor
2. Peripheral venous thrombosis
a. Spontaneous
b. Secondary to an antecedent stressor
3. Peripheral arterial thrombosis/thromboembolism (fatal or non-fatal)
c. Cerebrovascular – <i>Neurology</i>
1. Ischemic cerebrovascular stroke (fatal or non-fatal) with adequate documentation to subclassify as follows:
a. Large-artery atherosclerosis
b. Cardioembolism
c. Small-artery occlusion (lacune)
d. Other determined etiology
2. Ischemic cerebrovascular stroke (fatal or non-fatal) without adequate documentation to subclassify etiology
3. Hemorrhagic cerebrovascular stroke or hemorrhagic change (fatal or non-fatal)
4. Transient ischemic attack
5. Cerebrovascular venous thrombosis (fatal or non-fatal)
d. Non-Thromboembolic event

The SOP for this adjudication of cardiovascular events seem appropriate to FDA reviewers. The SOP includes a summary of the available data used by the

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committee and guidelines on the interpretation of cardiac data (e.g., how to interpret an elevated CPK-MB or an abnormal ECG).

Table 14. Listing of events referred for adjudication and adjudication results

Site/ Allocation Age/sex/ Treatment	Prior Hx/ CV risk factors	Low dose ASA	Adjudication
0064 4746 54M R	HTN	Yes	Y Non fatal MI
0126 2401 70F R	COPD, varicose ve	No	No Superficial thrombophlebitis
0193 5751 78F R	DM, CAD	No	Y Unstable angina
0200 3700 74M R	CAD, CABG	Yes	Y Sudden death
0212 1955 70F R	Atrial fib	No	No Aneurism. Subarachnoid hemorrh
0215 4378 72F R	1Lipid, CVA, ERT	Yes	No Non thromboembolic*
0357 0047 69M R	IBS, Peyronie's	No	Y Non fatal MI
0644 2176 79M R	1Lipid, HTN, PVD	Yes	Y Non fatal MI
0658 4049 71M R	HTN, CAD, Afib	No	Y Sudden death
0760 3253 72F R	HTN, hypothyroidism	Yes	No Non thromboembolic event
0810 6272 76M R	HTN, CAD	No	Y TIA. (w/Ventricular thrombus)
0821 5108 72M R	1Lipid, HTN,DM	No	Y Non fatal (Non Q wave) MI
0831 5382 70M R	HTN, angina	No	Y Non fatal MI
0002 9009 60M R	1Lipid, HTN, SyndrmX	Yes	No Non thromboembolic*
0016 9145 71F N	DM	No	Y TIA
0187 0665 60M N	-	No	Y Non fatal MI
0283 2182 58F N	? Hx of CVA, ERT	Yes	Y Ischemic CVA (no subclassif)*
0314 1477 70M N	CAD, HTN	No	No Non thromboembolic
0340 6823 67M N	CAD,HTN, DM	No	No Worsening CHF
0386 3155 77F N	1Lipid, HTN,DM	No	Y Ischemic CVA, small artery occlus*
0408 4129 54M N	CAD,CABG, HTN	Yes	Y Unstable angina
0443 1418 45F N	1Lipid, DM	No	Y Unstable angina
0449 4783 70F N	HTN, depression, ERT	No	Y Deep venous thrombosis
0462 1867 84F N	DM, postop per, ERT	No	Y Deep venous thrombosis**
0521 5761 58F N	HTN, ERT	No	Y Cerebral infarction
0580 6099 80M N	1Lipid, HTN	No	Y Ischemic CVA
0614 2792 74M N	HTN, prostatic ca.	No	Y Deep venous thrombosis
0702 6480 59F N	Cerebral hemorrh, ERT	No	Y Ischemic CVA
0774 3189 66F N	HTN, CAD, MI, ERT	No	Y Ischemic CVA
0065 5005 73F R	1Lipid, syst murmur	No	- Sudden death***

Source: Advantage CSR, CV adjudication package. R: rofecoxib. N: naproxen. CAD: coronary artery disease. HTN: hypertension, DM: diabetes mellitus. ERT: Estrogen replacement therapy. * FDA reviewers do not agree with adjudication. ** Occurred outside window determined by SOP and should have not been referred for adjudication. *** Not referred for adjudication.

Results of the adjudication:

Of the 29 cases evaluated by the CV adjudication committee (14 and 15 in the rofecoxib and naproxen groups respectively), seven were considered non-thromboembolic events, resulting in 9 and 12 adjudicated serious CV/thrombotic events.

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1.5.2.3 Review of serious CV events by FDA reviewers.

Reviewer's comment:

- *The referral for adjudication of two additional cases obtained from WAES - not reported by the investigators as cardiovascular thrombotic events - is of concern. One of these cases was adjudicated by the CV adjudication committee (102 462 1867, DVT on naproxen). However, a hand written note in the CRF for this case states that it is not known whether the patient ever took the assigned medication because she did not return the diary. As per the sponsor's June 22, 2001 correspondence, the event occurred more than 14 days after discontinuation of study drug. The SOP establishes a 14-day window period. Therefore, this case should have never been referred for adjudication.*
- *One cardiovascular death that should have been referred for adjudication was not referred because the term used by the investigator was not in the list of potential cardiovascular thrombotic events (hypertensive heart disease). This case was actually a case of sudden death (102 065- 5005, on rofecoxib). See Appendix 3a ("Terms eligible for adjudication").*
- *Post-hoc determination of the 'thrombotic' nature of some of these events may be difficult. Final interpretation of some data - often limited data - is by necessity, somewhat subjective. The Medical Officer assigned to this NDA reviewed all adjudication packages. For those cases in which the MO did not agree with the committee, adjudication packages were provided to the Divisions of Cardio-Renal and Neuropharm products. Two different FDA reviewers disagreed with the adjudication of four cases (0215 4378 211 and 0002 9009 308, on rofecoxib; and 0386 3155 209 and 0283 2182 222 on naproxen).*

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Table 15. Cases for which FDA reviewers disagree with the results of CV adjudication committee*.

Patient	Treatment	Adjudication Committee	CVT	FDA reviewers	CVT
0215 4378	Rofecoxib	Non-thromboembolic	N	Ischemic CVA	Y
0386 3155	Naproxen	Ischemic CVA, small artery occlusion	Y	Unable to adjudicate, Chorea of unknown etiology	N
0283 2182	Naproxen	Ischemic CVA	Y	Unable to adjudicate	N
0002 9009	Rofecoxib	Non-thromboembolic	N	Unstable angina	Y
0462 1867	Naproxen	DVT	Y **	Out of adjudication period	N
0065 5005	Rofecoxib	Death due to HTN	***	Sudden death	Y

Re-adjudication based on review of cases by reviewers from HFD-550, HFD-110 and HFD-120). CVT: serious cardiovascular thrombotic event. * Narratives are presented in Appendix 4. ** It should not have been referred for adjudication. *** It was not referred for adjudication.

Table 16. ADVANTAGE: Summary CV Thrombotic events as presented by the sponsor (adjudicated by CV Adjudication Committee) and FDA re-adjudicated events.

	Number of patients with CV Committee adjudicated serious CV-thrombotic events		Number of patients with FDA re-adjudicated serious CV-thrombotic events	
	Rofecoxib (N= 2785)	Naproxen (N= 2772)	Rofecoxib (N= 2785)	Naproxen (N= 2772)
	9	12	12	—
Cardiac	8	3	10	3
Sudden death	2	0	3	0
MI	5	1	5	1
Angina	1	2	2	2
Cerebrovascular	1	7	2	5
CVA	0	6	1	4
TIA	1	1	1	1
Peripheral	0	2	0	1
DVT/thromboflebit	0	2	0	1

There were **10 cardiac** events in the **rofecoxib** arm compared to **3** in the **naproxen** arm. Eight of the 10 cardiac events in the rofecoxib arm (5 MI, 2 sudden deaths, one angina) were in males; two additional cardiac events were in women. The cardiac cases in the naproxen arm were 2 unstable angina (1 man, 1 woman) and one MI (in a man with no prior CV history). Most of the cardiac events were in patients older than age 60, with known risks factors for CAD such as hypertension, diabetes, hyperlipidemia or prior history of CAD. Five of the ten cardiac events were in patients taking ASA. The number of events is small to allow interpretation regarding distribution of events in ASA users and non users.

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Reviewer's comment: A trend towards an excess of cardiac events in the rofecoxib 25 mg group in this 12 week study is noted. This observation is consistent with the pattern seen in VIGOR.

Of note, there were 2 and 5 cerebrovascular thrombotic events in the rofecoxib and naproxen arm, respectively, as adjudicated by the FDA Neuropharm reviewer. Two of the four CVA's were in patients taking hormonal replacement therapy. The reason for a lack of consistency between VIGOR and ADVANTAGE in regards to the risk of CVA's is unclear. It may be that cardiac and cerebrovascular events are different entities. However, the lack of power in the ADVANTAGE study may be an explanation.

A summary of the sponsor analyses of serious investigator reported CV events, adjudicated serious cardiovascular thrombotic and APTC (Anti Platelet Trialist's Collaboration) endpoints is presented in Appendix 3b. While the 95% CI's cross, the trend is consistent with FDA review for cardiac events.

The number of events is small, however, the finding is concerning because:

- 1. This is only a three month study.*
- 2. The dose of rofecoxib is 25 mg a day, the approved dose in OA (not twice the dose, as in VIGOR),*
- 3. The OA population is known to have lower cardiovascular risk than the RA population.*

The study is of extremely short duration to evaluate the cardiovascular effect of a drug in the prevention or increased risk of MI and stroke. Studies designed to evaluate cardiovascular outcomes are usually of 3-4 years duration.

1.5.3 NSAID-related events. The sponsor provided analyses of pre-specified NSAID-related adverse events similar to the ones that had been done in the VIGOR study.

1.5.3.1 NSAID-related adverse events related to edema, HTN and CHF

In this short-term, 12-week study, more patients had CHF related events and discontinued due to edema-related and HTN-related events in the rofecoxib 25 mg/day group than in the naproxen 1000 mg/day group.

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Table 17. ADVANTAGE study. NSAID-related adverse events related to edema¹, hypertension² and congestive heart failure³.

	Rofecoxib N= 2785		Naproxen N= 2772	
	n	%	n	%
Patients with Edema-related events	151	(5.42)	135	(4.87)
discontinued due to Edema related event	19	(0.68)	12	(0.43)
Patients with HTN related event	90	(3.23)	72	(2.59)
discontinued due to HTN related event	15	(0.54)	7	(0.25)
Patients with at least one CHF-related event	11	(0.39)	6	(0.22)

(Source Table 73, 74, 75 and appendix 4.1.57, Advantage CSR). Note: patients with two or more adverse events within a body system is counted only once within a body system. ¹

Includes terms such as edema, peripheral edema and lower extremity edema. ² Includes terms such as blood pressure increased, diastolic hypertension, systolic hypertension, labil hypertension, hypertension, uncontrolled hypertension.

Of the discontinuations due to edema-related events, only one patient (AN 1757, rofecoxib) was considered to have a serious event. The patient, a 72 year old male with history of HTN, DM, CAD, hyperlipidemia and CABG developed anasarca and was hospitalized approximately on treatment day #40. The physician indicated that the patient lost about 30 pounds of fluid with diuretic therapy during hospitalization.

CHF related events were defined as either congestive heart failure or left ventricular failure. Of the CHF related events, six were considered to be serious by the investigator: four in rofecoxib (AN 0085, 1082, 7389, 5108) and two in naproxen (6823 and 6398). Patient 5108 (a 72 year-old male) had a non-Q wave MI.

Reviewer's comment: The number of events is relatively small but despite the short duration of treatment and the fact that this is the 25 mg dose, the trend is consistent with findings in the VIGOR study.

1.5.3.2 Renal safety

There was one case of acute renal failure (AN 3105/777) in the naproxen arm, and one case of acute tubular necrosis (AN 1332/537 in the rofecoxib arm).

Twice the number of patients presented increased serum creatinine and BUN in the rofecoxib compared to the naproxen arm, respectively. Only one patient (in the naproxen arm) required discontinuation due to increased creatinine.

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Table 18 Renal related Laboratory AE experiences

	Rofecoxib N=2785 n (%)	Naproxen N=2772 n (%)
Blood urea nitrogen increased	15 (0.5)	7 (0.3)
Serum creatinine increased	21 (0.8)	10 (0.4)

1.5.3.3 Liver safety

There were two discontinuations due to liver-related clinical or laboratory adverse events, one in each treatment arm.

1.5.3.4 GI safety

The sponsor defined primary endpoint for this study was the cumulative incidence of discontinuations due to GI AE and abdominal pain at end of study endpoint.

The sponsor conducted a post-hoc analysis of PUBs (perforation, symptomatic ulcers and bleedings) similar to the analysis conducted for the VIGOR study. Of all the serious AE related to the digestive system, 6 and 12 events were referred to the GI adjudication committee from the rofecoxib and naproxen arms, respectively. Of those, 2 and 9 were confirmed PUBs by the adjudication committee. Of the confirmed PUBs, one was complicated in the rofecoxib group (the patient was taking ASA) and four were complicated in the naproxen group (one patient was taking ASA). The number of cases is small but the trend is consistent with the findings in VIGOR.

Reviewer's comment: The study succeeded in the sponsor's defined primary endpoint. However, this endpoint is not of clinical significance unless it is consistent with the overall safety profile, including cardiovascular events which are highly morbid. Only 7 of the 113 and 21 of the 142 discontinuations due to digestive symptoms were considered serious adverse events in the rofecoxib and naproxen arm respectively. Five of the 7 and 12 of the 21 serious digestive AE's required hospitalization in the rofecoxib and naproxen arm, respectively. Two of the seven and 3 of the 21 serious digestive AE's were on patients using concomitant aspirin in the rofecoxib and naproxen arm, respectively.

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2.0 Safety Update Report (SUR)

2.1 Overall Safety.

The Complete SUR was submitted to the Agency in July 12, 2001. This report includes the safety update of the extension studies in the original OA program (029, 058, 034 and 035), studies _____

_____ and 078, —and 091 (prevention of Alzheimer's). The SUR, however, does not contain the complete study reports for these studies.

Table 19. Studies included in Safety Update Report

Study/protocol #	Duration	Dose of rofecoxib	Rofecoxib	Patients randomized*	
				Placebo	Active control
Extension to Original NDA OA studies 034, 035	some patients up to >2 years	12.5 mg 25 mg 50 mg	415 (1) 475 78	-	409 Diclofenac (1)
Elderly— 058	Some patients up to 2 years	12.5 mg 25 mg	46 25	-	36 Nabumetone
New completed trials					
_____	Up to 65 weeks	25 mg	136	100	148 Ibuprofen (1)
Alzheimer's Disease— 091	Up to 479 days	25 mg	346 (14)	346 (8)	-
_____	6 weeks	25 mg 50 mg	53 49	59	-
_____	4 weeks	25 mg 50 mg	232 233	228	-
_____	6 weeks	12.5 mg	471 (1)	-	473 Naproxen
_____	6 weeks	12.5 mg	453 (1)	-	456 (Diclofenac/ misoprostol)
New ongoing trials					
Alzheimer's Disease— 078	Up to 1052 days	25 mg	721 (15)	729 (9)	-
_____	Up to 430 days	25 mg	381 (4)	376 (3)	-

* Patients randomized (Few patients were actually exposed to 2 years in the extension to the original NDA). The number of patients who died for any cause are in parenthesis. Source: modified from sponsor's table 1 of the SUR submitted 7/12/01 and data on deaths from individual studies.

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2.1.1 Deaths in the SUR.

There were a total of 58 deaths in the SUR. The number of deaths appears in parenthesis in Table 17.

Two deaths occurred during the extension period of the original NDA OA studies (029,034,035, 058):

- AN 8353, 64 M on rofecoxib 12.5 mg (post-op complication after CABG, day 607 of therapy)
- AN 5568, study 034: 75 F on diclofenac 150 mg, day 719 (post-op complication of hip replacement).

The total number of deaths in the entire original NDA and extensions was: 4 on rofecoxib 12.5 mg, one on rofecoxib 50 mg (in a patient with RA), one on naproxen, one on nabumetone and 9 deaths in the diclofenac group. For more detailed review the reader is referred to the MO review of NDA 21-042.

One death occurred in each of the remaining protocols submitted in this SUR:

- AN 0063, (protocol — a 60-year-old white woman with a history of hypertension and obesity was found dead in bed approximately 5 months into receiving ibuprofen 800 mg TID.
- AN 2301, (protocol — a 55 year old woman with history of chest pain and palpitations was found dead in bed 4 days after taking rofecoxib 12.5 mg a day. Autopsy showed coronary artery disease.
- AN 1659 (protocol — a 94-year-old man (AN 1659) on rofecoxib 12.5 mg for 15 days committed suicide by putting a plastic bag over his head.

Fifty-three of the 58 deaths were in the Alzheimer's studies: 33 on rofecoxib 25 mg and 20 on placebo (see below). Excluding the Alzheimer's studies, the number of deaths were small and do not raise additional safety concerns.

2.1.2 Serious AE's and Discontinuations due to AE's in the SUR

The pattern of SAE's and discontinuations due to AE's in the SUR was consistent with that observed in the original NDA. Of note, except the Alzheimer's studies most of these studies were of small size and short duration.

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2.2 Alzheimer's studies

Of all the data submitted in the SUR, the → studies for prevention of Alzheimer's disease potentially provide the most valuable information about long term exposure to rofecoxib 25 mg in comparison to placebo. However, limited safety data has been supplied from these studies. At the time of the submission of the SUR, study 091 had been completed; study 078 was ongoing and the data were not frozen by the cutoff date for the SUR (April 2001);

The SUR included listing of serious fatal and nonfatal AE's from the studies as well as adjudication packages of all serious potentially cardiovascular thrombotic events referred for adjudication to the CV adjudication committee. In subsequent submissions, at request of FDA reviewers, listing of discontinuations due to AE's and analyses of HTN, edema and CHF related events were provided.

Protocol 091 was a placebo-controlled, parallel-group, multicenter, 15-month double-blind study to evaluate the efficacy and safety of rofecoxib 25 mg to slow the progression of symptoms of Alzheimer's Disease. Patients of either gender who were ≥50 years of age with possible or probable Alzheimer's Disease were eligible to participate. Patients using NSAIDs for ≥7 days/month for the 2 months immediately prior to entry were not eligible. Patients were excluded if they were living in a nursing home or skilled nursing facility. Eligible patients were randomized to rofecoxib 25 mg or placebo for 12 months. This was followed by an additional 3-month treatment phase in which 90% of the patients initially assigned to rofecoxib were treated with placebo while the other patients remained on their initial treatment. Safety and tolerability were assessed at each visit (screening, Months 1, 3, 6, 9, 12, 13.5, and 15).

Protocol 078 This is a placebo-controlled, parallel-group, double-blind, multicenter study to evaluate the effects of rofecoxib 25 mg on the prevention of Alzheimer's Disease and cognitive decline in patients ≥65 years of age with mild cognitive impairment. Eligible patients were randomized to receive rofecoxib 25 mg or placebo for 2 years or until 220 cases of clinically diagnosed probable or possible Alzheimer's Disease are observed, whichever comes later. Safety and tolerability were to be assessed at all visits.

As per the sponsor's listings the demographic characteristics, co-morbid conditions and concomitant medications were similar in both treatment groups in each study. The mean age of these patients was 75 years and the number of patients included in each study per treatment arm was approximately 370 for study 091 approximately 700 for study 078.

The Alzheimer's studies specifically excluded patients at high cardiovascular risk. The following are some of the exclusion criteria used in the Alzheimer's studies (protocol 078):

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- 1 Patient with a history (within 2 years) or current evidence of major stroke, multiple lacunar infarcts or transient ischemic events.
- 2 Patient with a history of angina or congestive heart failure with symptoms at rest.
- 3 Patients with a history of myocardial infarction or coronary artery bypass grafting, angioplasty, or stent placement within 1 year prior to study start.
- 4 Patients taking the following medications:
 - Warfarin, heparin, ticlopidine.
 - NSAIDs (including salicylates or other aspirin-containing compounds) on a chronic basis (defined as ≥ 7 total days out of the last 30 days for 2 consecutive months prior to potential study entry).
 - Estrogen replacement therapies (excluding topical cream preparations)

At some point, patients on warfarin were made eligible for the study, provided that there was an increased frequency of monitoring of prothrombin time after initiation of blinded study therapy (amendment 78-02). In a later amendment, done after enrollment was complete, patients *who developed a need for cardio-protective doses of aspirin while in the trial* were permitted to use aspirin up to 100 mg/day(78-06).

Exposure

The following tables were presented by the sponsor in the SUR (July 12, 2001):

Table 20 (a, b and c). Patient exposure in Alzheimer's studies.

Number of Days on Therapy by Treatment—Protocol 091

	Treatment		
	Rofecoxib 25 mg/ Rofecoxib 25 mg	Rofecoxib 25 mg/ Placebo	Placebo/ Placebo
N	35	311	346
Mean	355.7	340.8	365.7
Range (min to max)	461 (2 to 463)	479 (1 to 480)	475 (1 to 476)

Number of Days on Therapy by Treatment—Protocol 078

	Treatment	
	Rofecoxib 25 mg	Placebo
N	721	729
Mean (SD)	500.16 (276.60)	549.50 (271.06)
Range (min to max)	1052 (0 to 1052)	1039 (0 to 1039)

Number of Days on Therapy by Treatment—Protocol —

	Treatment	
	Rofecoxib 25 mg	Placebo
N	381	376
Mean (SD)	155.96 (94.90)	161.84 (95.68)
Range (min to max)	349 (1 to 350)	429 (1 to 430)

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Reviewer's comment: As noted in the above tables, mean exposure to rofecoxib 25 mg in 346 patients in study 091 was 348.25 days (standard deviation not provided), with a range of one to 480 days. Mean exposure to rofecoxib 25 mg in 721 patients in study 078 was 500 days (± 276 days) with a range of 0 to 1052 days of treatment. Mean exposure to rofecoxib 25 mg in 381 patients in study — was 156 days (± 95 days) with a range of one to 350 days of treatment. The sponsor has not provided median time of exposure in each of the trials but has provided patient years at risk (see table below).

Table 21. Alzheimer's studies. Exposure to rofecoxib and placebo.

	Rofecoxib 25 mg		Placebo	
	Randomized	Pt. Years at risk	Randomized	Pt. Years at risk
091	346	301	346	366
078	721	996	729	1098
—	381	165	376	169
Total	1448	1461	1451	1634

Source: sponsor's table. SUR.

2.2.1 Overall Safety in the in Alzheimer's studies

2.2.1.1 Deaths

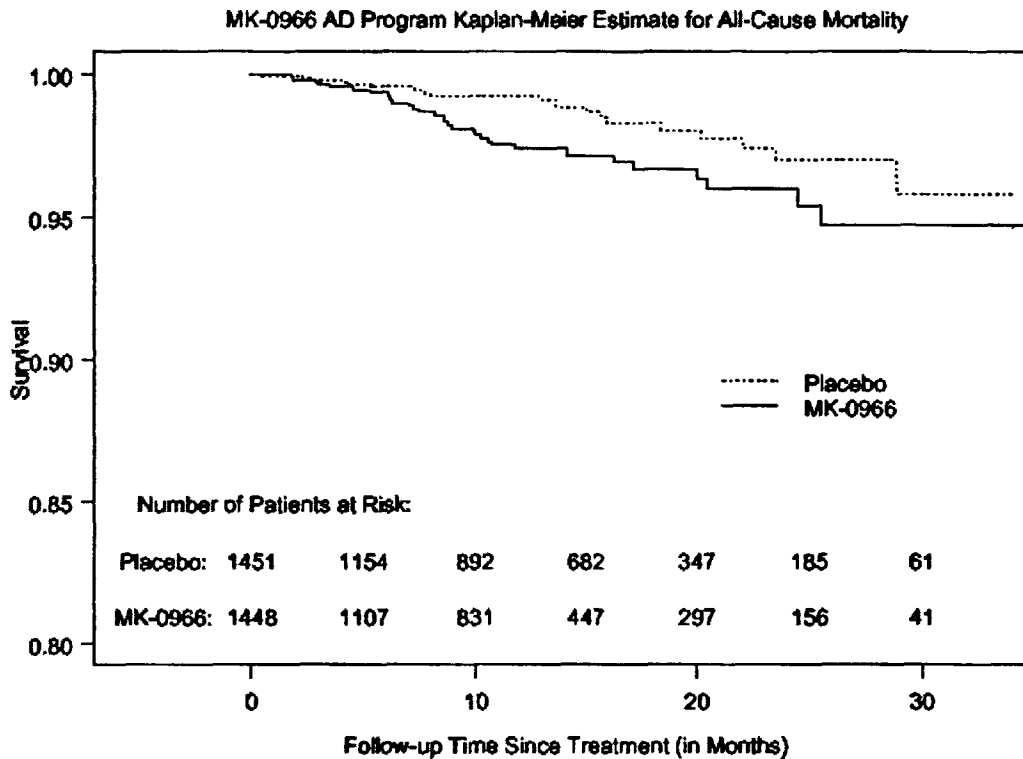
Pooled data from the — Alzheimer's studies showed that all cause mortality was higher in the rofecoxib 25 mg group, as compared to placebo (33 and 20 respectively). The p-value for the crude rate comparison between rofecoxib and placebo was 0.07. Of all deaths, 9 and 4 were confirmed as cardiovascular thrombotic by the CV adjudication committee.

Table 22. Deaths in Alzheimer's studies

Study #	Rofecoxib 25 mg N= 1448	Placebo N= 1451
	n/Pt Years of exposure	n/Pt Years of exposure
091	15/301	8/366
078	14/996	8/1098
—	4/165	3/169
	n (%)	n (%)
Total*	33 (2.3)	20 (1.4)
CVT ¹	8 (0.6)	4 (0.3)
Other ²	24 (1.7)	16 (1.1)

N= patients randomized. n= number of deaths. (%) crude rate. * P=0.007. ¹CVT: cardiovascular thrombotic death. Includes sudden death, myocardial infarction and ischemic cerebrovascular events confirmed as cardiovascular thrombotic by the CV adjudication Committee. ² Other: Includes death associated with malignancy, sepsis, trauma, CHF/pneumonia, pulmonary embolism, hemorrhagic stroke, unclassified cause of death.

Figure 1. Kaplan Meier estimates. All Cause Mortality in the Alzheimer's studies.



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The p-value for the logrank comparison between rofecoxib 25 mg and placebo = 0.026.
(Source of Kaplan Meier curve: provided by Sponsor on 11/12/01)

Reviewer's comment: Listing of the cause of death in all patients in Alzheimer's studies is presented in Appendix 6. Of note, the trend is consistent in study 091 and 078. Study 078 is still ongoing. The sponsor has reported that _____ (Information submitted 11/26/01).

2.2.1.2 Serious Adverse events in Alzheimer's studies.

Review of Serious Adverse events from the pooled Alzheimer's studies did not show major differences in all serious AE's, serious cardiovascular thrombotic events or serious digestive system events between rofecoxib and placebo.

Table 23. SUR. Summary of Serious AE's in Alzheimer's studies.

	Rofecoxib 25 mg N=1448 n (%)	Placebo N=1451 N (%)
Patients with at least one event	261 (18.0)	260 (17.9)
Body as a whole	70 (4.8)	55 (3.8)
CV AE's	77 (5.3)	82 (5.7)
Digestive AE's	40 (2.8)	32 (2.2)
Musculoskeletal	37 (2.6)	29 (2.0)
Skin and Appendices	20 (1.4)	42 (2.9)

Reviewer's comment:

There was a slightly higher number of serious digestive events in the rofecoxib group (2.8%) as compared to the placebo group (2.2%). These events included 2 gastric ulcers, 5 GI bleeding, 1 GI perforation, 2 hemorrhagic duodenal ulcer, 4 hemorrhagic gastric ulcer in the rofecoxib arm (N=12) and 1 GI bleeding, 2 GI perforation, 2 hemorrhagic gastric ulcer in the placebo arm (N=5) in the placebo arm. Of note, these are investigator reported terms, not confirmed events.

There was a slightly lower number of reported serious CV events in the rofecoxib group (5.3%) as compared to the placebo group (5.7%).

Discontinuations due to AE's were provided by the sponsor only for study 091. In this study of approximately 300 patients per treatment arm, there were differences in the digestive system (4.2 and 1.4% for rofecoxib and placebo respectively); respiratory system (2.3 and 0.3%) and urogenital system (1.9 and 0% had renal insufficiency) in the rofecoxib and placebo group, respectively.

2.2.2 Cardiovascular Safety in Alzheimer's studies

2.2.2.1 Cardiovascular thrombotic events referred for adjudication.

There was no substantial difference in the number of investigator reported serious cardiovascular potentially thrombotic events referred for adjudication between rofecoxib (n= 81) and placebo (n= 76). (Source: listing of serious CV thrombotic events referred for adjudication , submitted 9/6/01).

Reviewer's comment: By looking at the list of events referred for adjudication in the Alzheimer's studies it appears that the sponsor took a very conservative approach by referring all deaths, including terms such as "hepatic carcinoma" or "pulmonary fibrosis". However, even if only terms included in the original

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"Vascular SAE Terms Eligible for Case Adjudication" (Appendix 3a.) were included, the number of patients with events was not substantially different: 62 and 60, in the rofecoxib 25 and placebo groups, respectively. Of those, the sponsor reports that 22 and 30 were considered adjudicated APTC events in the rofecoxib and placebo groups, respectively.

The size of the database is relatively small to detect differences in cardiovascular safety. The — Alzheimer's studies all together had approximately 1500 patients on rofecoxib 25 mg and 1500 on placebo. As per calculations made by FDA statisticians and presented at the February 8, 2001 AAC meeting, based on the cumulative data of serious CV thrombotic events observed in VIGOR, at least 2500 patients would be needed in each treatment group to detect statistically significant differences between treatments, if they existed at the same rate as presented in VIGOR.

Of note, although the studies included an elderly population (mean age 75 years), patients with high cardiovascular risk such as those with a recent history of myocardial infarction and stroke, and patients taking estrogen replacement therapy were excluded from the Alzheimer's studies. Studies specifically designed to evaluate cardiovascular effects (e.g. protective effect of a drug in the risk of myocardial infarction or stroke) usually involve thousands of patients for several years.

The Alzheimer's studies showed a trend towards an excess of cardiovascular deaths in the rofecoxib 25 mg group: 8 and 4 confirmed cardiovascular thrombotic deaths, in the rofecoxib 25mg and placebo groups, respectively.

Data provided with the Alzheimer's studies do not cancel out the findings in the VIGOR and ADVANTAGE studies.

2.2.2.2 Hypertension-related, edema related and CHF related events

At the reviewer's request, the sponsor provided data on hypertension, edema and CHF related events in studies 091 — submitted 10/1/01). Data from study 078 were not available because this study is still ongoing and blinded.

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Table 24. SUR. Summary of HTN, edema and CHF-related events in Alzheimer's studies 091

	Rofecoxib 25 mg N= 726 n (%)	Placebo N= 722 N (%)
HTN-related	63 (8.7)	19 (2.6)
Edema-related	21 (2.9)	6 (0.8)
CHF-related	16 (2.2)	6 (0.8)

* Nine patients discontinued rofecoxib therapy due to the above AE's (3 in each category). One patient discontinued placebo due to a HTN- related event.

Reviewer's comment: consistent with prior databases, rofecoxib 25 mg is associated with higher incidence of hypertension, edema and CHF related events than placebo.

2.3 Short term placebo controlled studies in the SUR.

As part of the SUR, at the Agency's request, the sponsor provided data from five recently completed short term (4-6 weeks) studies that compared rofecoxib to placebo or other NSAIDs not included in the initial ADVANTAGE SUR. Three of these studies were placebo controlled (112, 116 and 905). In these studies, involving 456, 947 and 317 patients in the rofecoxib 12.5, rofecoxib 25 mg and placebo treatment arms respectively, four patients presented serious cardiovascular thrombotic events (one myocardial infarction and three coronary artery disease events), all in the rofecoxib 25 mg group.

The size and duration of these studies is too small to assess cardiovascular safety differences, however, the data suggest an increased risk in the rofecoxib 25 mg group as compared to placebo.

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3.0 Safety from the RA efficacy studies (21-042/s012).

This is the summary of a preliminary safety review of the RA efficacy application. A more detailed review is presented in a separate review (NDA 21-042/s012).

The protocols included in the RA efficacy application had a complicated design, with some patients switching treatments between parts. The RA safety database contains approximately 2000 patients exposed to rofecoxib (12.5, 25 and 50 mg); 550 patients exposed to naproxen and 1000 patients exposed to placebo. The bulk of the exposure was to 3 and 6 months of treatment. Approximately 1500 patients were randomized to rofecoxib 25 mg (n= 797) and 50 mg (n= 677) in 3-month placebo controlled studies. Approximately 180, 140 and 80 patients were exposed to rofecoxib 25mg, rofecoxib 50mg and naproxen 1000 mg respectively, for one year or more.

3.1 Overall safety in the RA application database

There were a total of six deaths: four on rofecoxib, one on naproxen and one on placebo. None of the deaths were considered by the investigator to be treatment related. None of them were cardiovascular deaths.

The pattern of adverse events, discontinuations due to adverse events, laboratory AE's and vital signs was consistent with data submitted in the original NDA submission.

3.2 Cardiovascular safety in the RA application database.

3.2.1 Investigator reported serious cardiovascular thrombotic events

The risk of developing serious cardiovascular thrombotic events with rofecoxib 50 mg in the RA application safety database is higher than with naproxen (2.6 vs. 1.5 per 100 patient years). The findings are consistent with the VIGOR study. The cardiovascular risk for the 25 mg dose (2.2 per 100 patient years) is also higher risk than naproxen. Risk comparisons to the rofecoxib 12.5 mg and placebo groups is inadequate because of the difference in exposure.

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Table 25. RA database. Summary of Investigator Reported Serious CV thrombotic events and Adjudicated events.(mostly 3-6 mo. studies, some patients exposed up to 3 years; placebo-controlled studies were only 3 months duration).

Treatment	Investigator reported serious CV thrombotic	Patient-years at risk *	Risk per 100 pt years	Adjudicated events	Risk per 100 pt years
Studies 096, 097, 098 and 103 (pivotal and endoscopic)					
Placebo	2	160	1.3	1	0.5
Vioxx 12.5	3	29	10.3	3	10.3
Vioxx 25	11	501	2.2	6	1.2
Vioxx 50	11	430	2.6	7	1.6
Naproxen	6	406	1.5	1	0.3

* patient-years at risk, provided by sponsor. Additionally, study 068 had 4, 7 and 5 investigator reported serious cv thrombotic events in the VIOXX 25, 50 and naproxen respectively, but only deaths were referred for adjudication, therefore, events from study 068 are not included in this analysis.

Reviewer's comment: Number of events is small. Finding is consistent with VIGOR. There are more CV thrombotic events in rofecoxib groups (12.5, 25 or 50 mg) as compared to naproxen. Interpretation of placebo data is difficult given the small number of patients and short exposure.

3.2.2 Edema-related AE's in RA application database.

The number of patients with edema-related events was higher in the rofecoxib 25 and 50 mg groups as compared to naproxen.

Table 26. Edema related events* (Source: sponsor's table 13 and 22 RA SUR).

	Placebo		Rofecox 25		Rofecox 50		Naproxen	
	n/N	%	n/N	%	n/N	%	n/N	%
Placebo controlled phase (12 weeks)	15/989	1.5	39/797	4.9	23/677	3.4	9/516	1.7
Long-term continuous (one-year data)	-		36/491	7.3	30/458	6.6	15/296	5.1

* Includes terms such as edema, fluid retention, lower extremity edema, peripheral edema. n= patients with events. N= patients randomized.

3.2.3 Hypertension-related AE's in the RA application database.

Hypertension related events were observed two to three times more often in each of the rofecoxib arms, as compared to the naproxen arm or placebo. A higher percentage of patients presented important increase of blood pressure and required concomitant

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medication in the rofecoxib arms compared to the naproxen arm. More patients discontinued due to HTN related events from each of the rofecoxib arms as compared to the naproxen arm. Of note, more patients had a prior history of hypertension in the rofecoxib 12.5 mg group.

Table 27. Summary of Hypertension related events in RA application* . (Source Table 13 and 22 RA SUR).

	Placebo		Rofecox 25		Rofecox 50		Naproxen	
	n/N	%	n/N	%	n/N	%	n/N	%
Placebo controlled phase (12 weeks)	22/989	2.2	49/797	6.1	43/677	6.4	10/516	1.9
Long-term continuous (one-year data)	-		59/491	12.0	71/458	15.5	16/296	5.4

* Includes terms such as blood pressure increased, diastolic hypertension, hypertension, uncontrolled hypertension. n= patients with events. N= patients randomized.

Of note, hypertension related events were two to three times more common in the rofecoxib groups as compared to the naproxen group.

3.2.4 CHF related events in RA application database

Three CHF related events occurred during the placebo controlled and long term therapy periods. All in the rofecoxib 50 mg group. Two additional cases occurred in the extension period, one in rofecoxib 25 mg and one in rofecoxib 50 mg. The number of CHF events is small to draw definitive conclusions but is consistent with VIGOR in which rofecoxib 50 mg was associated with higher risk of developing CHF related events than naproxen.

Table 28. Summary of CHF-related events (Source: Tables 13, 22 and 31, RA SUR)

	Placebo		Rofe 25		Rofe 50		Naproxen	
	n/N	%	n/N	%	n/N	%	n/N	%
Placebo controlled phase (12 weeks)	0/898	0	0/797	0	1/677	0.1	0/516	0
Long-term continuous (one-year)	-		0/491	0	2/458	0.4	0/296	0

* Includes pulmonary edema, congestive heart failure and cardiac failure. n= patients with events. N= patients randomized.

Reviewer's comment: In this database, rofecoxib 25 mg and 50 mg had higher incidence of cardiovascular thrombotic events, HTN-, edema- and CHF-related events compared to naproxen 1000 mg/day or placebo.

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B. Additional relevant data

Appendix 1. Narratives of Deaths

Patients allocated to Rofecoxib

065- 5005. Cause of death listed as "Hypertensive heart disease"

73 F, Hx HTN, hyperlipidemia, soft systolic murmur and hyokalemia. Not on ASA. Sept 11, 99 allocated to rofecoxib.

Oct 21 99 patient called son with c/o SOB When son arrived the patient was dead. Last contact with patient was on Oct 6, 99 when she indicated lack of efficacy. An autopsy was performed. Cause of death as per the coroner was "hypertensive heart disease and that the manner of death was "natural".

In the opinion of FDA reviewers' this is a case of sudden death. **THIS CASE HAD NOT BEEN REFERRED FOR ADJUDICATION TO THE CV ADJUDICATION COMMITTEE.**

200- 3700. Cause of death listed as MI. This case was adjudicated as Sudden Death.

74 M, hx angina, CAD, hyperchol, hypothy, DM, CABG (1964), GI bleeding. Concom therapy symvastatin, lisinopril, cyclobenzaprine, ASA, furosemide, levothyroxine, glyburide. In August 18, 99 allocated to rofecoxib. In Sept 29, 99 patient was thought to have suffered a massive MI. He expired at home. Autopsy not performed.

210- 4856. Cause of death: Glioblastoma multiforme

71 F, allocated to rofecoxib on Sept 29, 99. On Oct 8,99 patient had change in mental status. Oct 21 left side weakness and seizures. CT scan of head, large glioma in the frontal lobe. Patient expired Feb 2, 00.

658 – 4049. Cause of death listed as MI. Adjudicated as Sudden Death.

71 M, hx Afib, HTN, CAD. Conc; digoxin, diltiazem. No ASA.

Oct 15, 99 allocated to rofecoxib.

Patient attended visti 2 on Nov 22, 99 witout compaints. On Dec 15, 99 family informed the investigator that patient had been found dead in his home.

679- 3423. Cause of death listed as arterial rupture. Autopsy: Ruptured Aortic Aneurism.

75 F, no cv risk factors. Aug 24, 99 allocated to rofecoxib. August 25 reported pruritus and metallic taste. Reported she was under process of diagnostic testing for hematuria. On morning Oct 6, 99 patient was found dead in her kitchen. Autopsy showed ruptured aortic aneurism.

Patients allocated to Naproxen

059-1841 Cause of death malignant neoplasm: gallbladder carcinoma

56F allocated to naproxen Jul 2, 99. Patient reported she has been hospitalized in 9/6/99 for cholecystectomy and was diagnosed with gallbladder carcinoma. In Nov 2, 99 the patient expired.

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702-7154 Cause of death: malignang neoplasm: pancreatic carcinoma.

79F, hx of DM, HTN , anasarca, abdominal pain with constipation. In Oct 27, 99 allocated to naproxen. Jan 4, 200 increasing abdominal pain and distention. CT scan of abdomen showed large pancreatic mass 5.7 cm in the tail of the pancreas w/metastatic disease to the liver. Patient expired Jan 26, 2000.

777-3105 Cause of death: acute renal failure; respiratory failure; acidosis

74 F, hx of HTN, CHF, depression, cellulitis. Baseline labs K 5.4 mEq/L, BUN 49 mg/dl, Creatinine 1.5 g/dL, uric acid 11.4 mg/dL.

Augst 26, 99 patienn allocated to naproxen. Conc Verapamil, lisinopril, furosemide, fluoxetine.

Dec 5, 99 admitted to ER with respiratory distress, Cxray c/w CHF pneumonia or both. EKG IV conduction delay, sinus rhythm and first degree AV block. Creatinine of 4.8. UA WBC 5-7; RBC 10-12, bacteria 3+, mucus 3+. Patient diagnosied with acute renal failure. Echocardiogram showed enlarged right sided chambers with severe tricuspid regurgitation and pulmonary HTN and calcified mitral valve anulus. Patient deteriorated with elevated WBC count of 1911 K/L, HB 8.5 mg/dL, severe mixed metabolic and respiratory acidosis. Patient expired in Dec 7/99.

Last contact between patient and study coordinator was Oct 21, 99, at the 6 week visit. At that time the patient had a cold and had no complaints. Additional follow up stated that study therapy was dc on Oct 20,99 due to bronchitis.

In summary, this is a patient with history of HTN, CHF, tricuspid regurgitation and pulmonary hypertension and borderline renal function who started naproxen 500 mg bid on August 99. In November 99 she presented to the ER with acute renal failure, respiratory failure and acidosis, and died two days later. The investigator considered the episode not related to study medication. The cause of death in this patient is not clear. We might hypothesize that an NSAID worsened her borderline renal function and exacerbated her CHF. However, she may also have been septic. She did have an elevated WBC of 19 K but there are no available blood/urine/sputum cultures.

0831- 7114 Cause of death: Non-small cell carcinoma

78 M, hx glaucoma, doe, COPD peripheral vascular disoorder and smooker. Conc included ASA. Allocated Nov 19 99. Nov 23 99 rx disc due to nabdominal bloating. Dec 5, 99 patient hospitalized and diagnosed with metastatic adenocarcinoma of the lung and died on Feb 9, 2000.

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Appendix 2. Serious AE's that resulted in hospitalization

Fifty-three out of 68 (78%) patients with serious AE's in the rofecoxib group and 48 out of 72 (67%) patients with serious AE's in the naproxen group, required hospitalization.

Appendix 2. Hospitalizations (Source: Sponsor's Table 60, appendix 4.1.62)

	Rofecoxib 25 mg N= 2785		Naproxen 1000 mg N= 2772	
	n	%	n	%
Patients with at least one serious AE requiring hospitalization	53	(1.9%)	48	(1.7%)
Body as a whole	7	(0.3)	8	(0.3)
Cardiovascular system	18	(0.6)	13	(0.5)
Digestive system	5	(0.2)	12	(0.4)
Endocrine	1	(0.0)	1	(0.0)
Eyes, ears, nose and throat	-		1	(0.0)
Hemic and lymphatic	-		1	(0.0)
Hepatobiliary system	2	(0.1)	-	
Immune system	-		1	(0.0)
Musculoskeletal system	7	(0.3)	5	(0.2)
Nervous system	4	(0.1)	1	(0.0)
Psychiatric disorder	3	(0.1)	1	(0.0)
Respiratory system	6	(0.2)	3	(0.1)

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Appendix 3a.

Vascular SAE Terms (CRISP Broader Term)

Eligible for Case Adjudication

In the absence of other events identifying adverse experiences (most of which are likely to be serious adverse events), the following events (marked in strikethrough font) will have a low likelihood (in and of themselves) of being thromboembolic events. They may follow thromboembolic events, but alone do not represent such events.

acute myocardial infarction	cerebrovascular disorder
angina pectoris	congestive heart failure
anterior spinal artery obstruction	cor pulmonale
aortic atherosclerosis	coronary artery disease
aortic disorder	coronary artery occlusion
aortoiliac obstruction	coronary artery stenosis
arterial embolism	coronary vasospasm
arterial occlusion	coronary vessel surgery complication
arterial thrombosis	cyanosis
asystole	deep venous thrombosis
atherosclerosis	electrocardiographic abnormality
atrial fibrillation	electromechanical dissociation
atrial flutter	embolic stroke
basilar artery obstruction	embolism
brachial artery occlusion	endocardial disorder
bundle branch block	endocardial thrombus
cardiac aneurysm	extracranial artery obstruction
cardiac arrest	extradural hemorrhage
cardiac catheter complication	femoral artery occlusion
cardiac dyskinesia	gangrene
cardiac output low	idioventricular rhythm
cardiac stress test abnormality	iliac artery occlusion
cardiac thrombosis	incomplete left bundle branch block
cardiogenic shock	intermittent claudication
cardiomyopathy	intracranial hemorrhage
cardiovascular disorder	intracranial venous sinus phlebitis
cardiovascular hemodynamics abnormality	ischemic heart disease
carotid artery disorder	lacunar infarction
carotid artery obstruction	left bundle branch block
cerebellar artery obstruction	lower extremity arterial occlusion
cerebellar hemorrhage	lower extremity ischemia
cerebral artery obstruction	myocardial infarction
cerebral atherosclerosis	myocardial infarction complication
cerebral hypoxia	myocardial reinfarction
cerebral infarction	myocardial rupture
cerebral ischemia	non-Q-wave myocardial infarction
cerebral thrombosis	nonspecific ST-T change
cerebrovascular accident	old myocardial infarction

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papillary muscle disorder
~~peripheral atherosclerosis~~
peripheral ischemia
peripheral pulse absent
peripheral pulse decreased
peripheral vascular disorder
popliteal artery occlusion
~~pulmonary edema~~
pulmonary embolism
pulmonary infarction
pulmonary thrombosis
pulmonary vascular disease
pulmonary veno-occlusive disease
pulse absent
Q-wave abnormality
Q-wave myocardial infarction
QRS complex abnormality
shock
sinus thrombosis
small vessel disease
ST segment abnormality
ST segment depression
ST segment elevation
ST-T change compatible with ischemia
subclavian steal syndrome
sudden death
superior vena cava thrombosis
T-wave abnormality
T-wave flat
T-wave inversion

thromboembolic stroke
thromboembolism
thrombolysis
thrombophlebitis
thrombophlebitis migrans
thrombosis
thrombotic microangiopathy
transient ischemic attack
ulnar artery occlusion
unstable angina
upper extremity arterial occlusion
upper extremity ischemia
~~varicosity~~
vascular disorder
vascular graft occlusion
vascular insufficiency
vascular occlusion
vasospasm
venous compression
venous disorder
~~venous insufficiency~~
venous occlusion
venous thrombosis
ventricular fibrillation
ventricular flutter
ventricular tachycardia
ventricular thrombus
vertebral artery obstruction
vertebrobasilar insufficiency

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(Source: NDA 21-042/s007, Appendix 3.2.1).

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Appendix 3.b Sponsor's cardiovascular analyses.

Table Summary analysis of investigator reported serious cardiovascular thrombotic SAE's in ADVANTAGE.

Cohort	Treatment Group	N	Patients with Events	PYR ¹	Rates ²	Relative Risk ³	
						Estimate	95% CI
Total cohort	Rofecoxib	2785	14	639	2.19	1.06	(0.50, 2.26)
	Naproxen	2772	13	629	2.07		
Low dose aspirin user	Rofecoxib	352	5	81	6.18	5.08	(0.57, 240.4)
	Naproxen	367	1	82	1.22		

¹Patient-years at risk

²Per 100 PYR

³Relative risk of rofecoxib with respect to naproxen from unstratified Cox model where the number of cases is at least 11, otherwise relative risk is ratio of rates.

Source: sponsor's table 80. Appendix 4.1.80

Table. Summary of analysis of adjudicated thrombotic CV SAE in ADVANTAGE

Cohort	Treatment Group	N	Patients with Events	PYR ¹	Rates ²	Relative Risk ³	
						Estimate	95% CI
Total cohort	Rofecoxib	2785	9	640	1.41	0.74	(0.31, 1.75)
	Naproxen	2772	12	629	1.91		
Low dose aspirin user	Rofecoxib	352	3	81	3.71	3.05	(0.24, 159.9)
	Naproxen	367	1	82	1.22		

¹Patient-years at risk

²Per 100 PYR

³Relative risk of rofecoxib with respect to naproxen from unstratified Cox model where the number of cases is at least 11, otherwise relative risk is ratio of rates.

Source: sponsor's table 78 Appendix 4.1.78

Table Summary of analysis of APTC endpoints in ADVANTAGE

Cohort	Treatment Group	N	Patients with Events	PYR ¹	Rates ²	Relative Risk ³	
						Estimate	95% CI
Total cohort	Rofecoxib	2785	10	640	1.56	1.41	(0.54, 3.69)
	Naproxen	2772	7	629	1.11		
Low dose aspirin user	Rofecoxib	352	3	81	3.71		
	Naproxen	367	0	82	0.00		

¹Patient-years at risk

²Per 100 PYR

³Relative risk of rofecoxib with respect to naproxen from unstratified Cox model where the number of cases is at least 11, otherwise relative risk is ratio of rates.

Source: sponsor's table 79 Appendix 4.1.79

APTC: Anti Platelet Trialist Collaboration endpoints: CV death, non-fatal MI, non-fatal stroke. (Difference with "Adjudication thrombotic endpoints": Includes hemorrhagic stroke. Excludes angina, TIA and peripheral arterial and venous events).

Appendix 4. Narratives of patients for which FDA reviewers did not agree with sponsor's adjudication.

102 462 1867 - 84 year old woman with history of DM, osteoporosis, hypothyroidism and partial gastrectomy allocated to receive naproxen in July 29 1999. Concomitant therapy included estrogens, insulin and synthroid. On August 30, 99 she was hospitalized for severe back pain with a vertebral compression fracture. She was transferred to a rehab center in 9/3/99. In 9/10/99 she developed leg edema, was diagnosed with DVT and started anticoagulation. Subsequently developed colon hemorrhage (9/15/99) secondary to anticoagulation. The case report form originally had 8/16/99 as the stop date for taking the study medication. The date was later corrected to 7/9/99. There is a hand written note stating that it is not known whether the patient ever took the assigned medication because she did not return the diary. As per the sponsor's June 22, 2001 correspondence, the event occurred more than 14 days after discontinuation of study drug.

102 065- 5005. 73 year old woman with history of HTN, hyperlipidemia, soft systolic murmur and hyokalemia. Allocated to rofecoxib in Sept 11, 1999. In Oct 21, 1999 patient called son with c/o SOB. When son arrived the patient was dead. Last contact with patient at the study site had been in Oct 6, 1999 when she indicated lack of efficacy. An autopsy was performed. Cause of death as per the coroner was "hypertensive heart disease" and that the manner of death was "natural". This is actually a sudden death.

102 0215 4378 211: 72 year old female with history of hyperlipidemia, CVA and carotid endarterectomy. Concomitant meds: ASA, clopidogrel, prevastatin, estradiol. Patient was randomized in Oct 6, 1999. In Oct 31, 1999 patient developed numbness and tingling similar to her prior CVA. She went to surgery. A thrombus of the right carotid artery with severe stenosis was found intraoperatively. This case was considered "non-thromboembolic" by the CV adjudication committee. The Division of Neuropharm products considered this case as an "ischemic cerebrovascular accident". This patient was on rofecoxib 25 mg.

102 0386 3155 209: 77 year old female with history of HTN, hyperlipidemia, hypothyroidism, DM and dyspepsia. Concomitant therapy included omeprazole, atenolol, nifedipine, atrovastatin, levothyroxine, clopidogrel temazepam, rosiglitazone, potassium and furosemide. Patient was randomized in June 18, 1999. In Aug 5, 1999, a physician reported that the patient developed abnormal head and arm movements diagnosed as dyskinesia of unknown cause. Study therapy was discontinued. Follow up information from a physician indicated that the diagnosis was changed to possible small CVA. In August 9, 1999 an MRI showed lacunar type infarctions bilaterally in the caudate region; an old hemorrhagic infarct in the right putamen and bilateral thalamic lacuna with no definitive evidence of acute infarct. Patient recovered within 5 days with "adjustment" of medications and haloperidol. This event was adjudicated as "ischemic with small artery occlusion". Acute choreic symptoms can rarely be caused by a lacunar infarct. Acute symptoms are most likely related to metabolic imbalances, e.g. hyperglycemia, which the patient had, exceeding 450 mg/dL during her hospital stay. The Division of Neuropharm products considered this case as "Unable to adjudicate. Acute chorea of unknown etiology". This patient was on naproxen.

102 0283 2182 222 58 year old female with history of cardiac murmur, allergic cough and intermittent anemia. Patient family history includes development of TIA's with symptoms including dysphagia. Patient was randomized in July 21, 1999. Concomitant medications

Clinical Review Section

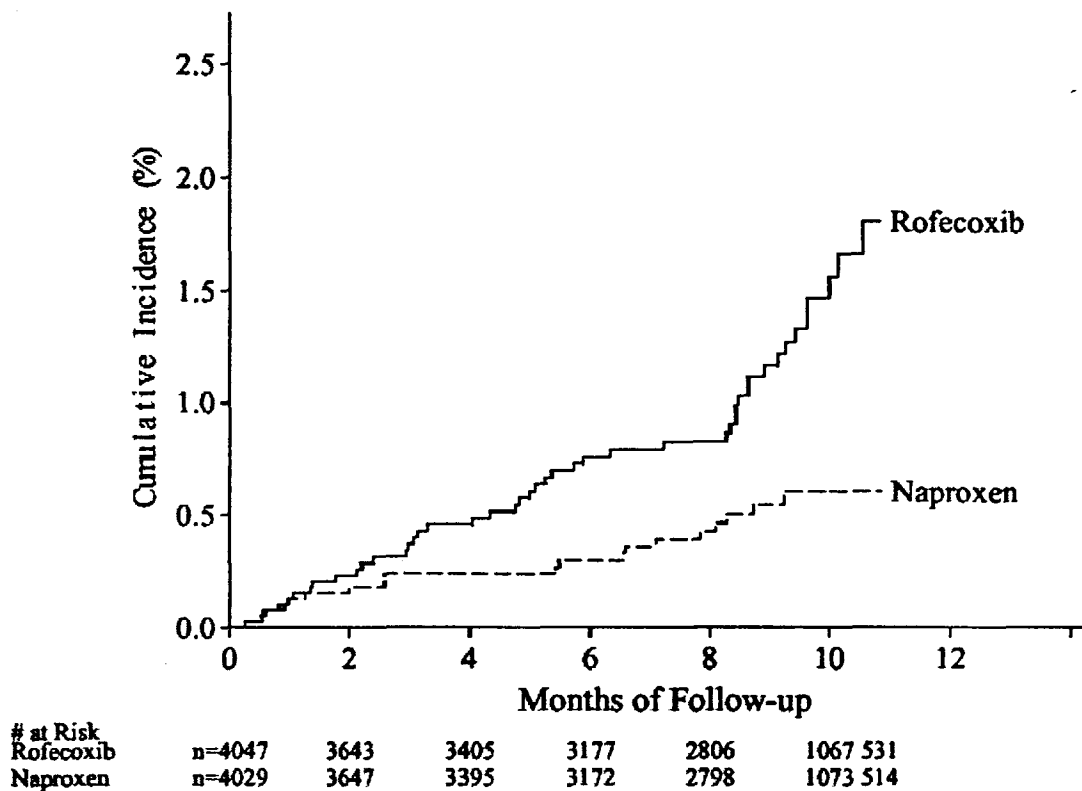
included conjugated estrogenic hormone w/medroxyprogesterone and famciclovir. Documentation for this case is poor and confusing. In August 5, 1999 for no particular reason, she was started on baby ASA "for general health". Apparently, starting August 16, 1999 the patient had one or two neurologic episodes characterized by speech difficulty and left side weakness in the context of severe headache. In September 11, 1999 ASA was increased from baby ASA to 235 mg/day. Between September 23, 1999 and October 12, 1999 she had another transient episode of dysphagia for 24 hours. The adjudication package and narrative provided by the sponsor is unclear as to when exactly those TIA's occurred. The patient did not look for medical attention at the time of the events. Additionally, recurrent episodes of weakness appeared to precede study entry. Neurologic examination did not confirm left side weakness. Head CT, carotid ultrasound and echocardiogram (September 24-27) were all normal. The patient completed the trial in October 12, 1999. The CV adjudication committee adjudicated these events as "ischemic CV stroke". The Division of Neuropharm products considered this case to be "Unable to adjudicate". The patient was on naproxen.

903 0002 9009 308. 60 year old male with a history of osteoarthritis, obesity, HTN, and hypercholesterolemia. His PMH is significant for a history of a subendocardial MI in 1990. A coronary angiogram in 1991 revealed 'normal coronary arteries with a bit slow flow'. He had several normal stress tests in the past. He was diagnosed with Syndrome X. The patient was randomized and took study drug for approximately 80 days before the onset of chest pain leading to hospitalization. Chest pain lasted 2.5 hours (6 pm to 8:30 pm), was associated with nausea and not affected by 6 nitroglycerin tablets. ECGs on admissions showed no ischemic changes. Per the hospital discharge summary, MI was excluded *despite an elevated CPK-MB (5 mcg/ml on admission, normals 0 to 3 mcg/ml) and troponin (0.8, normals 0-0.4)*. After an echocardiogram and a stress test that was reported as negative ('without steadily ground for ischemia') the patient's pain was attributed to Syndrome X and discharged. The event was adjudicated as not an APTC endpoint per the CV adjudication committee. Because of the persistently elevated CPK-MB and troponin, in the presence of chest pain accompanied by nausea and chest pressure, in patient with prior history of CAD (subendocardial MI) the division of Cardio-Renal products considered this event likely to be due to a cardiac thrombotic event. The patient was on rofecoxib.

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Appendix 5. a. Cardiovascular safety in VIGOR study

Confirmed (Adjudicated) Thrombotic CV serious adverse experiences in the VIGOR study. Time to event plot (all patients randomized). RR=2.37 for rofecoxib relative to naproxen (p=0.001).



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Appendix 5.b. VIGOR study. Confirmed or “adjudicated” thrombotic cardiovascular serious adverse experiences in all patients randomized and in subgroups of patients identified retrospectively by the sponsor as patients who may have or may have not benefited from low dose ASA.

	N	Patients with events (%)	PYR ¹	Rates ²	Relative risk ³		
					Estimate	95%CI	p
All patients randomized							
Rofecoxib	4047	45 (1.1%)	2697	1.67	2.37	1.39 – 4.06	0.0016
Naproxen	4029	19 (0.5%)	2698	0.70			
Potential candidate for low dose ASA ⁵							
Rofecoxib	170	15 (8.8%)	105	14.29	4.89	1.41 - 16.88	0.0122
Naproxen	151	3 (2.0 %)	102	2.94			
Not candidate for low dose ASA							
Rofecoxib	3877	30 (0.8%)	2592	1.16	1.88	1.03 – 3.45	0.041
Naproxen	3838	16 (0.4%)	2596	0.62			

¹ Patient-years at risk. ² Per 100 patients years. ⁴ Relative risk of rofecoxib with respect to naproxen. ⁵ Patients with past medical history of cerebrovascular accident, transient ischemic attack, myocardial infarction, unstable angina, stable angina, coronary artery bypass surgery or percutaneous coronary intervention. (Source: modified from sponsor’s Table 9 of the safety update, Estimate calculated by Dr. Qian Li, FDA statistician).

VIGOR study. Myocardial Infarctions. Subgroup analyses by sponsor’s retrospective identification of patients who may have benefited from low dose ASA.

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Appendix 5.c. VIGOR. Investigator reported and adjudicated (CV committee confirmed) serious CV thrombotic events. (Source: NDA 21-042/007, SUR 10/3/01 submission).

	Number of patients with Investigator reported serious CV thrombotic events		Number of patients with CV Committee confirmed serious CV-thrombotic events	
	Rofecoxib (N=)	Naproxen (N=)	Rofecoxib (N=)	Naproxen (N=)
	64	32	47	20
CV death	7	7	6	6
Fatal acute MI	3	4	2	-
Fatal hemorrhagic stroke	-	-	1	1
Fatal Ischemic stroke	2	1	-	1
Sudden cardiac death	2	-	3	4
Intracranial hemorrhage	-	2	-	-
Cardiac events (fatal & nonfatal)	36	19	28	10
MI	23	8	20	4
Angina	6	7	3	4
Vent fib	1	-	5	3
Cardiac arrest	1	-	-	-
Coronary art disease/occlusion	2	3	-	-
Ischemic heart dz.	2	1	-	-
Cerebrovascular (fatal& nonfatal)	20	11	13	9
CVA	15	6	-	-
Hemorrhagic stroke	-	-	2	1
Ischemic cerebr. vasc stroke	-	3	9	8
TIA	2	-	2	-
Carotid artery obstruction	1	-	-	-
Cerebrovascular disorder	1	2	-	-
Intracranial hemorrhage	1	2	-	-
Peripheral	8	2	6	1
Arterial thrombosis	1	1	1	-
Venous thrombosis	5	1	5	1
Peripheral vascular disorder	1	-	-	-
Arterial embolism	1	-	-	-

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Appendix 6. a. Deaths in Alzheimer's protocol # 091

Rofecoxib 25 mg (14)

- *AN 332 - Cardiac arrest , fatal MI: 78 M, 1 day off drug, relative day 328
- *AN 601 - Sudden death: 75 M, 1 day off drug, relative day 228
- *AN 831 - CVA: 85 W M, relative day 296
- AN 915 - CHF and pneumonia: 79 F, relative day 260
- AN 964 - Dizziness, CVA, respiratory failure: 86 F, taking conjugated estrogens, developed CVA approx. 1 month into the study. Drug stopped. Patient died on relative day 58. Not adjudicated due to insufficient data.
- AN 3 - Pneumonia: 91 F day 260
- AN 42 - Endocarditis/pneumonia: 83 M, day 303
- AN 282 - Lung cancer: 71 F, day 165
- AN 376 - Burn/fungemia/anuria: 62 M, day 173
- AN 382 - Lung and brain malignant neoplasm: 77 F, relative day 123
- AN 542 - Hypercalcemia, acute renal failure: 82 F 7 days off drug; relative day 188
- AN 691 - Esophageal malignant neoplasm: 86 M, relative day onset 390
- AN 835 - Interstitial lung disease, pneumonia, cardiac arrest: 82 M, day 269
- AN 891 - Fever, sepsis. 88 F, day 76

Placebo (8)

- *AN 784 - Sudden death: 70 F, day 458
- AN 394 - Metastatic neoplasm, unknown primary: 68 M, day 452
- AN 613 - COPD/ aspiration pneumonia: 88 M, day 131
- AN 664 - Alzheimer's/Pneumonia: 84 M, day 417
- AN 827 - Intracranial hemorrhage: 74 M, day 70 (CV non-thromboembolic)
- AN 830 - Acute myelogenous leukemia: 79 F, day 99
- AN 832 - Pneumonia, cardiac arrest secondary to aspiration/sepsis: 82 M, day 461
- AN 956 - Ruptured aortic aneurysm: 81 M, day 16 (CV non-thromboembolic)

* Case confirmed by Cardiovascular Adjudication Committee

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Appendix 6b. Deaths in Alzheimer's protocol # 078 (from WAES database)

Rofecoxib 25 mg (15)

*AN 248 – Sudden death, unknown cause of death: 71 F, day 747
*AN 359 - Sudden death, hypertension: 68 M, day 624
*AN 737 - Sudden death, cardiac arrest: 84 M, day 185
*AN 799 - Sudden death, cardiac arrest: 85 M, day 312
*AN1025 – Acute Myocardial infarction: 83 M, day 138
AN 205 - Postop complication, hip fracture, pulmonary embolism: 85 M, day 496 (CV)
AN 583 - Pulmonary embolism/ pancreatic ca: 70 M, day 357 (CV)
AN 352 - Hemorrhagic duodenal ulcer; small cell ca; 67 M, day 322
AN 762 - Metastatic prostate ca, renal failure; 87 M, day 707
AN 821 - Head trauma: 85 M, day 271
AN 935 - Trauma: 75 M, day 106
AN 1097 – Electric shock: 69 M, day 248
AN 1453 – Chest trauma: 83 M, day 611
AN1453 – Bacterial sepsis, acute myelogenous leukemia: 80 M, day 53

Placebo (8):

*AN 1256 – Sudden death: 82 F, day 674
*AN 1378 – Sudden death: 74 M, day 392
AN 539 - Hypertension: 72 M, day 243 (CV).
AN 264 – Colon ca: 76 M, day 430
AN 294 - malignant melanoma: 77 M, day 556
AN 308 - myelogenous leukemia, pneumonia, acute renal failure: 85 M, day 407
AN 1144 - Pancreatic carcinoma: 94 F, day 469
AN 1350 - bladder carcinoma: 79 F, day 708
AN 1547 - metastatic neoplasm of unknown origin: 82 M, day 96

Deaths in Alzheimer's protocol —

Rofecoxib (4)

AN 125 - GI (large intestine) perforation: 86 F, day 214
AN 466 - Hip fracture, dyspnea (PE?): 86 M, day 190 NOT REFERRED x adjudication
AN 532 - Hemorrhagic CVA (confirmed)
AN 743 - Intracranial hemorrhage (confirmed)

Placebo (3)

*AN 661 - Myocardial infarction, related to major stressor: meningitis: 77 M, day 236
AN 257 - Trauma: 79 M, day 82
AM 635 - Lymphoma, GI bleeding, sepsis: 78 F, day 216

* Confirmed as cardiovascular thrombotic event by CV Adjudication Committee. Of note, mean exposure in this study was 5 months only.

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Lawrence Goldkind
12/8/01 11:49:14 AM
MEDICAL OFFICER

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Correction to Table 10 and Table 16 of MO review of Complete Response to Approvable letter of April 6, 2001 (date of review: 11/28/01).

Table 16. ADVANTAGE: Summary CV Thrombotic events as presented by the sponsor (adjudicated by CV Adjudication Committee) and FDA re-adjudicated events.

	Number of patients with CV Committee adjudicated serious CV-thrombotic events		Number of patients with FDA re-adjudicated serious CV-thrombotic events	
	Rofecoxib (N= 2785)	Naproxen (N= 2772)	Rofecoxib (N= 2785)	Naproxen (N= 2772)
	9	12	12	1 9
Cardiac	8	3	10	3
Sudden death	2	0	3	0
MI	5	1	5	1
Angina	1	2	2	2
Cerebrovascular	1	7	2	5
CVA	0	6	1	4
TIA	1	1	1	1
Peripheral	0	2	0	1
DVT/thromboflebit	0	2	0	1

Table 10. Serious AE's by ASA use (events with incidence 0.5%), as reported by investigators.

	Rofecoxib N= 2785		Naproxen N= 2772	
	Non ASA* users	Low dose ASA	Non ASA users	Low dose ASA
	N= 2422	N= 352	N= 2398	N= 367
	n (%)	n (%)	n (%)	n (%)
Patients with at least one event in any body system	54 (2.2)	14 (4.0)	59 (2.5)	12 (3.3)
Cardiovascular	16 (0.7)	7 (2.0)	1 15 (0.6)	2 (0.5)
Digestive system	5 (0.2)	2 (0.6)	18 (0.8)	3 (0.8)

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FOOD AND DRUG ADMINISTRATION
DIVISION OF ANTI-INFLAMMATORY, ANALGESIC AND OPHTHALMIC DRUG
PRODUCTS -- HFD-550

Medical Officer Review

VIOXX
(Rofecoxib)

NDA 21-042 (capsules) and NDA 21-052 (oral solution)
S 007 (Gastrointestinal Safety)

Submission date (letter):	June 29, 2000
Submission type:	supplement NDA
End of Review date:	March 30, 2001
Reviewer:	Maria Lourdes Villalba, MD.
Drug name:	VIOXX (Rofecoxib)
Applicant:	Merck Research Laboratories
Pharmacologic category:	NSAID (COX-2 inhibitor)
Proposed indications:	Management of acute pain, dysmenorrhea and signs and symptoms of osteoarthritis.
Dosage form and route:	Oral capsule, 12.5, 25 mg and 50 mg Oral solution 12.5 mg/5ml and 25 mg/5ml
Project Manager:	Sandra Folkendt
Related Reviews:	Stats: Qian Li, Ph.D; GI safety: Lawrence Goldkind, M.D.; CV safety: Shari Targum, M.D. Original NDA 21-042 reviews. IND — (Rofecoxib). OPDRA safety reviews

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HFD-550/MO/Goldkind

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Lawrence Goldkind, M.D., Team Leader,
DAIAODP

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<p>Appendix 1. Financial Disclosure Appendix 2. VIGOR. Narratives of deaths Appendix 3. Summary of adjudicated serious thromboembolic events in selected subgroups of patients in VIGOR. Appendix 4. Discontinuations due to adverse events Appendix 5. Edema-related and HTN-related events in original NDA 21-042. Appendix 6. Thrombotic events in original NDA 21-042.</p>	

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Executive summary

1. Recommendations for Regulatory Action: Approvable.

In order to adequately interpret the cardiovascular and overall safety results in the VIGOR study and provide adequate labeling information, review of the complete report of study 102 (ADVANTAGE) is necessary. A review of the safety database from the ADVANTAGE trial will allow for further global safety assessment of rofecoxib at currently labeled chronic doses and in subjects requiring cardioprotective doses of aspirin.

The applicant should be informed that labeling changes cannot be made until review of safety database from the ADVANTAGE study is complete.

2. Summary of findings

2.1 Data sources:

NDA 20-042/052 supplement 007 included three new studies (088c, 085, 090), and two studies from the original NDA (study 058 and 069). Additionally, data submitted to — and post-marketing data were reviewed.

Study 088c (VIGOR) was a 8,000-patient study comparing rofecoxib 50 mg daily to naproxen 500 mg BID (median exposure was 9 months). The study excluded patients who were taking or were candidates for taking low dose aspirin (ASA). Studies 085, 090 and 058 were 6-week efficacy studies of rofecoxib 12.5 and 25 mg daily that allowed concomitant use of low dose ASA. Study 069 was a pooled analysis of GI events in the entire phase II/III osteoarthritis program. Except for study 058, all studies included in study 069 also excluded the use of low dose ASA.

As part of the — (rofecoxib), the applicant conducted a 12-week 5,500-patient safety study (# 102 or "ADVANTAGE"), comparing rofecoxib 25 mg daily and naproxen 500 mg BID in a population that did not exclude the use of low dose ASA. This study was completed in March 2000, however, the complete study report has not been submitted to the FDA for review. The FDA medical reviewers have requested that the complete report of study 102, be submitted for review.

2.2 Conclusions

2.2.1. GI safety

1. The sponsor demonstrated a statistically significant reduction in PUBs and complicated PUB's, associated with the use of rofecoxib compared to naproxen in

this population of patients not considered by their physicians to have an indication for cardiovascular prophylaxis with low dose ASA.

The cumulative incidence of PUB's was 1.8% and 3.9% (median exposure of 9 months) for rofecoxib and naproxen, respectively. Of note, this cumulative rate is very close to the 2 – 4% rate presented in the WARNING section of the NSAID template for patients treated for one year. The cumulative incidence of *complicated* PUB's was 0.5 % and 1.2 % in the rofecoxib and naproxen groups, respectively.

The target population for anti-inflammatory drugs includes a substantial number of patients who will need low dose ASA for cardiovascular prophylaxis. Adequate data on the safety of the chronic co-use of rofecoxib and low dose ASA do not exist. Studies that allowed the concomitant use of rofecoxib and low dose ASA were of short duration and not powered to detect differences in serious GI and CV events. Therefore, the exclusion of patients using low dose ASA is a serious limitation to the generalizability of the findings of the VIGOR study.

Different NSAIDs are associated with different risk of developing serious GI events. The only non-selective NSAID comparator included in this study was naproxen. Therefore, claims of GI superiority could be only be made in comparison to naproxen, not to all NSAIDs.

2. The post-marketing safety profile of rofecoxib is similar to other NSAIDs, including the risk of GI bleeding. From May 1999 to October 2000, the FDA post-marketing AER system received 37 unduplicated reports of death due to gastrointestinal complications associated with the use of rofecoxib. Despite a substantial risk reduction compared to naproxen in the VIGOR study, the risk of serious GI complications associated with rofecoxib is still a concern. Risk factors associated with serious GI complications are similar to those associated with conventional NSAIDs: age, prior history of ulcer disease, concomitant use of ASA, coumadin or other antiplatelet agents, and corticosteroids.
3. Data provided by the sponsor do not support removal of the NSAID class GI WARNING section from the VIOXX label.

2.2.2. Cardiovascular safety

1. **The cumulative rate for serious CV/thrombotic events was 1.8% (n= 45) and 0.6% (n= 19) in the rofecoxib and naproxen groups respectively over the 9-month period. The relative risk of developing serious CV/thrombotic events was more than twice in the rofecoxib group as compared to the naproxen group (RR= 2.37; 95% C.I 1.39, 4.06; p= 0.0016, based on risk per 100 patient years). The difference was mainly due to the difference in the number of myocardial infarction (MI): 20 in rofecoxib and 4 in naproxen (crude rate 0.5 % and 0.1% for rofecoxib and naproxen, respectively) (RR= 5.0; 95% C.I. 1.72, 14.3, based on risk per 100 patient years).**